# (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property International Bureau Organization



(43) International Publication Date 31 December 2003 (31.12.2003)

(10) International Publication Number WO 2004/001042 PCT

(21) International Application Number: PCT/DK2003/000417 C12N 15/10 (51) International Patent Classification7:

AZ BA, BB, GB, BR, BY, BZ, CA, Cli, CA, CO, CR, CU, CC, CR, CO, CR, CO, CR, CO, CR, CO, CR, CO, CR, CR, CD, CZ, DB, DR, DM, DZ, BC, EB, ES, FI, GB, GD, GB, GH, GM, HR, HU, DLI, NI, SI, PK, KI, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MX, NO, NO, PH, PI, PI, RO, RU, SC, SD, SB, SG, SK, SI, TI, TM, TY, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, PEDERSEN, Henrik [DK/DK]; Frodesvej 24, DK-2880 Bagsværd (DK). Ē

20 June 2003 (20.06.2003) (22) International Filing Date:

English (25) Filing Language:

English US OF US 20 June 2002 (20.06.2002) 20 June 2002 (20.06.2002) (26) Publication Language: PCT/DK02/00419 Priority Data: 10/175,539 8

PA 2002 01953

3

(71) Applicant (for all designated States except US): NUEVO-LUTION A/S [DK/DK]; Rønnegade 8, 5, DK-2100 19 December 2002 (19.12.2002) 20 December 2002 (20.12.2002) København Ø (DK). 60/434,664

Inventors/Appilicants (for US only): FRESKGÅRD, Per-Ola [SE/SE]; Ringvägen 7, S-235 93 Vellinge (SE). Inventors; and <u>8</u>8

without international search report and to be republished upon receipt of that report Published:

W. Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Emasian patent (AM, AZ, BY, KG, KZ, MD, RU, TI, TM), European patent (KH, BH, BG, CH, CY, CZ, DB, DK, BE, ES, FI, FK, GB, GR, HU, LB, TI, LU, MC, NL, PT, RO, SR, SI, SK, TR), OAPI patent (BF, BI, CF, CG, CI, CM, GA, GW, ML, MR, NE, SN, TD, TG).

5

For two-letter codes and other abbreviations, refer to the "Guid-ance Notes on Codes and Abbreviations" appearing at the begin-ning of each regular issue of the PCT Gazette.

WO 2004/001042

PCT/DK2003/000417

## Microarrays displaying encoded molecules

### Technical Field of the Invention

been annealed to a complex so as to present at spatially defined spots en-The present invention relates to an oligonucleotide microarray which has coded molecules

### Background

9

genotype of an Individual. Information on the expression of the genes may be Microarrays have found a wide acceptance in various analysis concepts. Miknowledge on a disease state, hormone action, infection etc. The presence croarrays can be used to profile gene patterns yielding Information on the accomplished using messenger RNA (mRNA) samples in order to obtain or absence of a particular SNP may also be verified using Microamays.

proteins. As spots on the microarray comprise different sequences of nucleothereof. The mRNA-protein fusion is formed by allowing a ribosome to transate an mRNA provided at the 3' end with puromycln and linking the formed in US 60207,446 B1 it has been suggested to use a microarray to present protein to said mRNA at the termination of the translation. By annealing a nixture of RNA-protein fusions to a microarray it is possible to display the ides and the RNA sequences anneal sequence specifically to the probes, polypeptides linked to the mRNA which is responsible for the formation he proteins are presented at spatial defined areas of the microarray.

8

22

not limited to the reaction products of the 20 naturally occurring amino acids, which allow for a higher diversity of the presented molecule and the possibilnvention, it is small molecules which are presented and in another aspect it cording to an object of the present invention it is desired to expand the type of molecules which can be presented on a microarray. In one aspect of the The prior art is restricted to the presenting of proteins on a microarray. Acis unnatural polymers that are presented. Notably, the present invention is

ဓ

CONFIRMATION COPY

(57) Abstract: Disclosed is a microarray comprising a plurality of single stranded nucleic acid probes immobilized in discrete areas of a solid support, said probes being hybridised to a library of complexes, wherein each complex comprises an encoded molecule and a template which codes for said molecule, said template comprising a number of codons which codes for chemical entities which upon reaction form a reaction product which at least partly form part of the encoded molecule.

001 001 0 (54) Thie: MICROARRAYS DISPLAYING ENCODED MOLECULIES

2 (57) Abstract: Disclosed is a mirrarmy communicate a classificate of the stand o

7V

conditions, such as high temperature, extreme pH and in media containing lty of forming robust and stable molecules that can be treated under harsh

### Summary of the Invention

ა

stranded nucleic acld probes Immobilized in discrete areas of a solid support, chemical entities which upon reaction form a reaction product which at least The present invention relates to microarray comprising a plurality of single plex comprises an encoded molecule and a template which codes for said sald probes being hybridlsed to a library of complexes, wherein each commolecule, said template comprising a number of codons which codes for partly form part of the encoded molecule.

9

ent single stranded oligonucleotide probes immobilized in discrete areas of a Suitably, an ollgonucleotide microarray is a device having a plurality of differ-The term microarray generally refers to an ordered array of microscopic elements on a planar substrate. Commercially available standard oligonucleosolid support. The discrete areas comprising immobilized single-stranded tide microarrays may be used to prepare the microarray of the Invention. oligonucleotides may be referred to as spots for short.

5

ಜ

low. Preferred oligonucleotides are capable of forming a specific hybridisation The oligonucleotides immobilised in the spots may comprise any oligomer of nucleotides known in the art and In particular the nucleotides described bewith a complementing oligonucleotide.

22

be used. To adapt to the scanners usually used in the art, the solid support is support is preferable flat, that is, the solid support has even parallel surfaces The solid support is preferable dimensional to add precision to the manufacusually of the dimension of a traditional 1  $\times$  3-in microscope slide. The solid turing and detection steps. Any specific dimension of the spotting area may over a local region. The solid support should be uniform in the sense that

ജ

support is a glass plate, a silicon or silicon-glass plate (e.g. a microchip), or a contribute any gain or loss of signal in the detection step. Usually, the solid irregularities preferably are avoid in the bulk of the support as well as in the surface coating or treatment. Preferably, the solid support is durable, i.e. a processed microarray should loose less than 10% of the annealed oligonucleotides over the assay duration, and inert, i.e. the solid support does not polymer plate.

ics and readily detection. The centre-to-centre spacing of the spots is sultable at least one other spot comprises a different nucleic acid probe. The distance Each spot on the solid support comprises the same nucleic acid probe, while A, and preferably between 10 and 50 A to allow for optimized reaction kinetbetween each Immobilized oligonucleotide on the spot is suitably 10 to 100 constant and In the range from 20µm to 1000µm and more preferred be-9

tween 50µm and 500µm.

5

The ollgonudeotides immobilized on the solid support can be prepared in any convenient way. Usually, either a delivery approach or a synthesis approach is used. According to the delivery approach the oligonucleotides are syntheslsed, e.g. using the phosphoramidate method, and subsequent printed on

- of an aldehyde surface on the solld support and reacting with an oligonucleoent attachment, and (lii) the covalent attachment of an oligonucleotide carryzation of preformed oligonucleotides may be performed utilizing any of a variety of attachment chemistries, such as (i) the formation of aminosilanes on the solid support, where the oligonucleotides are immobilized. The immobilia glass support and attaching the oligonucleotides thereto, (ii) the formation 01/04129. The synthesis approach employs in situ synthesis of the oligonuide comprising an amine, typically an aliphatic amine linker to form a covaing an anthraquinone to a polymer solid support as disclosed in WO 32 ន
- final oligonucleotide eventually is formed. Usually, a method employing photo cleotides on the solld support using repeated addition of nucleotides until the activation and masking is used. ജ

Each of the complexes hybridised to the microarray comprises an encoded molecule and a template which codes for the encoded molecule. Usually, the template comprises a nucleic acid, such as an ollgonucleotide. The oligonucleotide can contain any of the nucleotides mentioned below.

2

It is preferred that the template is divided into coding regions or codons which codes for specific chemical entities. A codon is a sequence of nucleotides or a single nucleotide. The nucleotides are usually amplifiable and the nucleobases are selected from the natural nucleobases (adenine, guanine, uracil, thymine, and cytosine) and the backbone is selected from DNA and RNA, preferably DNA.

으

The codon may be a single nucleotide. In the generation of a library, this will allow for the incorporation of four different chemical entities into the encoded molecule. However, to obtain a higher diversity a codon in certain embodiments preferrably comprises at least two and more preferred at least three nucleotides. Theoretically, this will provide for 4² and 4³, respectively, different chemical entities. The codons will usually not comprise more than 100 nucleotides. It is preferred to have codons with a sequence of 3 to 30 nucleo-

5

20

The template will in general have at least two codons, which are arranged in sequence, I.e. next to each other. Each of the codons may be separated by a spacer group. Depending on the encoded molecule formed, the template may comprise further codons, such as 3, 4, 5, or more codons. Each of the further codons may be separated by a suitable spacer group. Preferably, all or at least a majority of the codons of the template are arranged in sequence and each of the codons is separated from a neighbouring codon by a spacer group. Alternatively, codons on the template may be designed with overlapping sequences.

22

PCT/DK2003/000417

Generally, it is preferred to have more than two codons on the template to allow for the synthesis of more diverse template-directed molecules. In a preferred aspect of the invention the number of codons of the template is 2 to 100. Still more preferred is templates comprising 3 to 20 codons.

- The spacer sequence may serve various purposes. In one setup of the Invention, the spacer group identifies the position of a codon. Usually, the spacer group either upstream or downstream of a codon comprises information which allows determination of the position of the codon. The spacer group may also or in addition provide for a region of high affinity. The high affinity region will ensure that the hybridisation of the template with the anti-codon will occur in frame. Moreover, the spacer sequence may adjust the ennealing temperature to a desired level.
- A spacer sequence with high affinity can be provided by incorporation of one or more nucleobases forming three hydrogen bonds to a cognate nucleobase. An example of a nucleobase having this property is guanine. Alternatively, or in addition, the spacer sequence may be subjected to back bone modification. Several back bone modifications provides for higher affinity, such as 2'-O-methyl substitution of the ribose molety, peptide nucleic acids (PNA), and 2'-4' O-methylene cyclisation of the ribose molety, also referred to as LNA (Locked Nucleic Acid).

5

2

The template may comprise flanking regions. The flanking region can encompasses a signal group, such a flourophor or a radio active group, to allow a direct detection of the presence of the complex. The flanking regions can also serve as priming sites for an amplification reaction, such as PCR. The template may in certain embodiments comprise an affinity region having the property of being able to hybridise to a building block.

22

It is to be understood that when the term template is used in the present description and claims, the template may be in the sense or the anti-sense format, i.e. the template part of the complex can be a sequence of codons

which actually codes for the molecule or can be a sequence complementary

more chemical bond attaching the encoded molecule to the template in order product may be post-modified to obtain the encoded molecule displayed on reacted with each other and/or a scaffold molecule. Optionally, this reaction the microarray. The post-modification may involve the cleavage of one or The encoded molecule is formed by a variety of reactants which have more efficiently to display the encoded molecule.

ß

connection to another reactive group positioned on a chemical entity, thereby generating an addition to the original scaffold. A second chemical entity may comprises an amine group a connection between these can be mediated by The formation of an encoded molecule generally starts by a scaffold, I.e. a entities maybe involved in the formation of the final reaction product. The reactive group Incorporated by the first chemical entity. Further chemical example, if the nascent encoded molecule and the chemical entity both chemical unit having one or more reactive groups capable of forming a formation of a connection between the chemical entity and the nascent react with a reactive group also appearing on the original scaffold or a encoded molecule maybe be mediated by a bridging molecule. As an a dicarboxylic acid.

5

2

ឧ

15

suitable linking moiety. Furthermore, the encoded molecule may be linked to the template through a cleavable linker to release the encoded molecule at a The encoded molecule may be attached directly the template or through a point in time selected by the experimenter.

22

prior to the participation in the formation of the reaction product leading the generally comprises an anti-codon, in some embodiment the building block eliminations of the encoded molecule may be attached to a building block final encoded molecule. Besides the chemical entity, the building block The chemical entities that are precursors for structural additions or

ဓ

ဗ္က

also comprise an affinity region providing for affinity towards the nascent complex.

molecule by a building block, which further comprises an anticodon. The antihowever, it is important that a correspondence is maintained in the complex transfer of genetic information and chemical entity may occur in any order, Thus, the chemical entities are suitably mediated to the nascent encoded codon serves the function of transferring the genetic Information of the building block in conjunction with the transfer of a chemical entity. The ည

ribosomes or enzymes having similar activity. The chemical entitles may be The chemical entities are preferably reacted without enzymatic interaction. Votably, the reaction of the chemical entities is preferably not mediated by reacted with each other or a scaffold having a reciplent reactive group. 9

anti-codon is transferred by specific hybridisation to a codon on the template. the nascent complex are to anneal an oligonucleotide complementary to the anti-codon and attach this ollgonucleotide to the complex, e.g. by Ilgation. A According to certain aspects of the invention the genetic information of the Other methods for transferring the genetic information of the anti-codon to

codon to the nascent complex using a polymerase and a mixture of dNTPs. still further method Involves transferring the genetic information of the anti-

ន

The chemical entity of the building block serves the function of being a pre-

25

derstood that not necessarily all the atoms of the original chemical entity is to cursor for the structural entity eventually incorporated into the encoded molecule. Therefore, when it in the present application with claims is stated that a chemical entity is transferred to a nascent encoded molecule it is to be unnolecule. Especially, the cleavage resulting in the release of the entity may chemical entity can be changed when it appears on the nascent encoded be found in the eventually formed encoded molecule. Also, as a consequence of the reactions involved in the connection, the structure of the

The chemical entity of the building block comprises at least one reactive group capable of participating in a reaction which results in a connection between the chemical entity of the building block and another chemical entity or a scaffold associated with the nascent complex. The connection is facilitated by one or more reactive groups of the chemical entity. The number of reactive groups which appear on the chemical entity is suitably one to ten. A

ß

building block featuring only one reactive group is used I.a. in the end positions of polymers or scaffolds, whereas building blocks having two reactive groups are suitable for the formation of the body part of a polymer or scaffolds capable of being reacted further. One, two or more reactive groups intended for the formation of connections, are typically present on scaffolds. A scaffold is a core structure, which forms the basis for the creation of multiple variants. The variant forms of the scaffold are typically formed through reaction of reactive groups of the scaffold with reactive groups of other building blocks, optionally mediated by fill-in groups or catalysts, under the creation of a connection between the entities. The chemical entities to be connected to the scaffold may contain one, two or several reactive groups able to form

The reactive group of the building block may be capable of forming a direct connection to a reactive group of the nascent complex or the reactive group of the building block may be capable of forming a connection to a reactive group of the nascent complex through a bridging fill-in group. It is to be understood that not all the atoms of a reactive group are necessarily maintained in the connection formed. Rather, the reactive groups are to be regarded as precursors for the structure of the connection. Fig. 5 shows examples of various reactive groups and the corresponding connection formed.

25

೫

WO 2004/001042

PCT/DK2003/000417

თ

The subsequent cleavage step to release the chemical entity from the building block can be performed in any appropriate way. In an aspect of the Invention the cleavage involves usage of a reagent or and enzyme. The cleavage results in a transfer of the chemical entity to the nascent encoded molecule

or in a transfer of the nascent encoded molecule to the chemical entity of the building block. In some cases it may be advantageous to introduce new chemical groups as a consequence of linker cleavage. The new chemical groups may be used for further reaction in a subsequent cycle, either directly or after having been activated. In other cases it is desirable that no trace of the linker remains after the cleavage. Fig. 6 shows examples of conditions for linkers between the building block and the chemical entity to be cleaved.

In another aspect, the connection and the deavage is conducted as a simultaneous reaction, i.e. either the chemical entity of the building block or the

र

nascent encoded molecule is a leaving group of the reaction. In general, it is preferred to design the system such that the connection and the cleavage occur simultaneously because this will reduce the number of steps and the complexity. The simultaneous connection and cleavage can also be designed such that either no trace of the Ilnker remains or such that a new chemical

group for further reaction is introduced, as described above. In fig. 4 exemplary reactive groups leading to transfer of a chemical entity to the entitles harbouring one of the reactive groups are shown.

ಜ

It is important for the method according to the invention that at least one linkage remains intact between the encoded molecule and the template. In case the method essentially involves the transfer of chemical entities to a scaffold or an evolving polymer, the eventually scaffolded molecule or the polymer may be attached with a selectively cleavable linker. The selectively cleavable linker is designed such that it is not cleaved under conditions which result in

22

30 a transfer of the chemical entity to the nascent encoded molecule.

tion of pyrimidines. The nucleotide may be distanced from the reactive group of the chemical entity by a spacer molety. The spacer may be designed such cleobase is used for attachment of the chemical entity, the attachment point phosphor of the internucleoside linkage or at the nucleobase. When the nuis usually at the 7 position of the purines or 7-deaza-purins or at the 5 posithat the conformational spaced sampled by the reactive group is optimized The attachment of the chemical entity can be at any entity available for attachment, e.g. the chemical entity can be attached to a nucleobase or the back bone. In general, it is preferred to attach the chemical entity at the for a reaction with the reactive group of the nascent encoded molecule.

building block:template hybrids to ensure that the anti-codons have been anobtaining annealing temperatures in a specific range for all or some of the nealed to the template before the chemical entities are connected to each The design of building blocks comprising the anti-codon may be aimed at other through a chemical reaction.

5

9

within a certain range to obtain a condition at which all or at least the majority The templates are preferably designed to have an annealing temperature of complexes can be annealed to the probes of the array through the templates.

8

The complexes for the library can be prepared in accordance with a variety of methods. Examples of these methods are depicted below and generally

described In PCT/DK 02/00419. 23

oligonucleotides are provided. Subsequently a primer is annealed to the each unnatural nucleotides as building blocks. Initially, a plurality of template A first embodiment is based on the use of a polymerase to incorporate derivatives which have appended chemical entities. Subsequent to or template and a polymerase is extending the primer using nucleotide

ဓ္က

simultaneously with the incorporation of the nucleotide derivatives, the chemical entitles are reacted to form a reaction product. Several possible reaction approaches for the chemical entitles are apparent.

- groups, the chemical entities can be attached to adjacent chemical entities by a reaction of these reactive groups. Exemplary of the reactive groups are polymerised. In the event the chemical entities each carry two reactive First, the nucleotide derivatives can be incorporated and subsequently amine and carboxylic acid, which upon reaction form an amide bond.
- nterspaced reactive group to obtain a linkage to the chemical entity, e.g. by approach is the use of a reactive group between a chemical entity and the oridging molety. Exemplary of this approach is the linking of two chemical entities each bearing an amine group by a bi-carboxylic acid. Yet another nucleotide building block, such an ester or a thioester group. An adjacent building block having a reactive group such as an amine may cleave the Adjacent chemical entities can also be linked together using a linking or 9 5

an amide linking group.

- hybridisation of building blocks to a template and reaction of chemical entitles approach comprises that a plurality of templates are contacted with a plurality of building blocks, wherein each building block comprises an anti-codon and A second embodiment for obtainment of complexes pertains to the use of attached to the building blocks in order to obtain a reaction product. This a chemical entity. The anti-codons are designed to have a recognition 8
  - sequence, i.e. a codon, on the template. Subsequent to the annealing of the anti-codons and the codons to each other a reaction of the chemical entities are effected to obtain a reaction product. 22

reaction of the reactive group of the chemical entity may be effected at any The template may be associated with a scaffold. Building blocks bringing chemical entities in may be added sequentially or simultaneously and a time after the annealing of the building blocks to the template. ဓ္တ

PCT/DK2003/000417

12

A third embodiment for the generation of a complex includes chemical or enzymatical ligation of building blocks when these are lined up on a template. Initially a plurality of templates are provided, each having one or more

- codons. The templates are contacted with building blocks comprlsing anticodons linked to chemical entities. The two or more anti-codons annealed on a template are subsequently ligated to each other and a reaction of the chemical entities is effected to obtain a reaction product.
- A fourth embodiment makes use of the extension by a polymerase of an affinity sequence of the nascent complex to transfer the anti-codon of a building block to the nascent complex. The method implies that a nascent complex comprising a scaffold and an affinity region, which may and may not contain coding regions. A building block comprising a region complementary to the affinity section is subsequently annealed to the nascent complex and the anti-codon region of the building block is transferred to the nascent complex by a polymerase. The transfer of the chemical entity may be transferred prior to, simultaneously with or subsequent to the transfer of the anti-codon.

5

9

After or simultaneously with the formation of the reaction product some of the linkers to the template may be cleaved, however at least one linker must be maintained to provide for the complex.

8

The library of the complexes may be added to the oligonucleotide microarray under hybridisation conditions in order for each template to anneal to a cognate probe on the microarray. The hybridisation conditions may be appropriately adjusted by a person skilled in the art taking into account the number and kind of nucleobases that participate in the formation of the

ജ

22

WO 2004/001042

PCT/DK2003/000417

೮

It is within the capability of the skilled person in the art to construct the desired design of an oligonucleotide. When a specific annealing temperature is desired it is a standard procedure to suggest appropriate compositions of nucleic acid monomers and the length thereof. The construction of an appropriate design may be assisted by software, such as Vector NTI Suite or the

http://www.nwfsc.noaa.gov/protocols/oligoTMcalc.html.

public database at the internet address

S

The conditions which allow specific hybridisation of the templates and probes are influenced by a number of factors including temperature, salt

- concentration, type of buffer, and acidity. It is within the capabilities of the person skilled in the art to select appropriate conditions to ensure that the contacting between the templates and the probes are performed at hybridisation conditions. The temperature at which two single stranded oligonucleotides forms a duplex is referred to as the annealing temperature
- or the melting temperature. The melting curve is usually not sharp indicating that the annealing occurs over a temperature range. The second derivative of the melting curve is used herein to indicate the annealing temperature.

The array according to the present Invention may have many uses. One aspect of the present Invention relates to methods for detecting the presence or absence of, and/or measuring the amount of target molecules in a sample, wherein the method employs complexes comprising encoded molecules, which is attached in an array system. The target molecule in the sample which has an affinity towards the encoded molecule immobilised on the mi-

- croarray will bind to the encoded molecule this making the detection or measuring possible. In the event a variety of target molecules are present in the sample and a variety of encoded molecules are presented, multiple determinations are rendered possible.
- 30 The invention described herein provides arrays that can measure different amounts at the protein level without the use of proteins or peptides as detection molecules. The template-displaying molecule technology could be used

S

deposition of thousands of different functional molecules onto different loca-The template-encoded motecule complexes can be directly applied to a prespotted DNA and hybridised the probe, optionally using an adapter oligonuother possibility is that the synthesis could be performed directly on the precoated template using a polymerase and the nucleotide analogues. Making cleotide, which is complementary to the probe as well as the template. Anaddressable microarrays with this technology will lead to high-throughput ions of a chip. The overall principal is shown in figure 1.

9

5

2

bated with a microarray displaying encoded molecules. By this approach, the Ordered display of encoded molecules is a powerful tool for the Identification process in which valldated targets are used in order to identify suitable bindlabelled due to binding of the target. Related targets can be compared in order to identify molecules with high specificity for a specific subtype by analyidentity of encoded molecules that are able to bind to a target molecule are format, a target is detectably labelled, e.g. with a fluorescent dye, and incucules. The above specific format may find application in the drug discovery determined from the location of the spots on the microarray that becomes sls the binding pattem of these related targets on identical arrays of moleof previous unknown target-encoded molecule interaction. In one specific

23

5

parts, namely a nucleobase moiety, and a backbone. The back bone may in ome cases be subdivided into a sugar moiety and an intemudeoside linker. The nucleotides used in the present invention may be linked together in an oligonucleotide. Each nucleotide monomer is normally composed of two ß

tydroxy-5-methyl-4-triazolopyridine, isocytosine, isoguanine, inosine and the bases are adenine, guanine, thymine, cytosine, 5-methylcytosine, and uracil, \forall ,N⁴-ethanocytosin, \N<sup>6</sup>, \N<sup>6</sup>-ethano-2,6-dlamino-purine, 5-methylcytosine, 5-(C3-C8)-alkynylcytosine, 5-fluorouracil, 5-bromouracil, pseudoisocytosine, 2diaminopurine, 8-oxo-N<sup>6</sup>-methyladenine, 7-deazaxanthine, 7-deazaguanine, No. 5,432,272. The term "nucleobase" is intended to cover these examples deobases are adenine, guanine, thymine, cytosine, uracii, purine, xanthine, as well as analogues and tautomers thereof. Especially interesting nucleohe nucleobase molety may be selected among naturally occurring nucleooases as well as non-naturally occurring nucleobases. Thus, "nucleobase" non-naturally occurring" nucleobases described in Benner et al., U.S. Pat neterocyclic analogues and tautomers thereof. Illustrative examples of nuncludes not only the known purine and pyrimidine hetero-cycles, but also 5

9

Examples of suitable specific pairs of nucleobases are shown below:

which are considered as the naturally occurring nucleobases in relation to

ಜ

therapeutic and diagnostic application in humans.

22

PCT/DK2003/000417

Re-H: Unacido Coposino Coposin

othetic Base Pal

thetic purine bases pairring with natural pyrim

Suitable examples of backbone units are shown below (B denotes a nucleobase):

ß

6

The sugar molety of the backbone is suitably a pentose but may be the appropriate part of an PNA or a six-member ring. Suitable examples of possible pentoses include ribose, 2'-deoxyribose, 2'-O-methyl-ribose, 2'-flour-ribose, and 2'-4'-O-methylene-ribose (LNA). Suitably the nucleobase is attached to the 1' position of the pentose entity.

An internucleoside linker connects the 3' end of preceding monomer to a 5' end of a succeeding monomer when the sugar molety of the backbone is a pentose, like ribose or 2-deoxyribose. The Internucleoside linkage may be the natural occurring phospodiester linkage or a derivative thereof. Examples of such derivatives include phosphorothioate, methylphosphonate,

PCT/DK2003/000417

phosphoramidate, phosphotriester, and phosphodithioate. Furthermore, the intermucleoside linker can be any of a number of non-phosphorous-containing linkers known in the art.

Preferred nucleic acid monomers include naturally occurring nucleosides forming part of the DNA as well as the RNA family connected through phosphodiester linkages. The members of the DNA family include decoxyadenosine, deoxyguanosine, deoxythymidine, and deoxycytidine. The members of the RNA family include adenosine, guanosine, uridine, cytidine, and inosine. Inosine is a non-specific pairing nucleoside and may be used as universal base as discussed above because inosine can pair nearly isoenergetically with A, T, and C.

9

S

### Brief Description of the Figures

15

- Fig. 1 shows an array of displayed encoded molecules.

  Fig. 2 shows the synthesis of an array of encoded molecules.

  Fig. 3 shows the selection to obtain encoded molecules for array analysis

  Fig. 4 shows reaction types allowing simultaneous reaction and cleavage, i.e. transfer of chemical entities to a nascent encoded molecule.
- 20 Fig. 5 shows pairs of reactive groups and the resulting bond.
  Fig. 6 shows various cleavable chemical bonds, the products obtained and the agents necessary for the cleavage to occur.

## Detailed description of the figures

25

Fig. 1 shows an encoded molecule chip. A DNA library is spotted in array format on a suitable surface. A library of complexes comprising single-stranded template DNA is added and allowed to hybridise to the complement DNA strand. This allows site-specific immobilization of the encoded molecules. The site-specific immobilization is controlled by the codons of the

30 complex.

WO 2004/001042

PCT/DK2003/000417

6

Fig 2 shows the hybridization of encoded molecules mediated by the codon composition in each template molecule. According to a specific use, a biological sample containing target molecules is added and non-bound material is washed off. The final step is the detection of bound material in each single spot. In this example, the target molecule is labelled for detection.

တ

Fig. 3 shows the selection of an Initial encoded library against a target. The enriched encoded molecules are hybridized on the array due the precise combination of codons in the template molecule. Finally, 'target molecule(s) are allowed to bind to the arrayed oligonucleotides and detected.

2

Fig. 4 shows different classes of reactions which mediate transfer of a chemical entity from a building block to another entity, or to an anchorage point. For simplicity reasons only, the receiving entity is shown as a building block; it is to be understood that the receiving entity can be covalently attached to the template as well. The reactions have been grouped into three different classes: Nucleophilic substitutions, addition-elimination reactions, and transition metal catalyzed reactions.

5

(A) Reaction of nucleophiles with carbonyls. As a result of the nucleophilic substitution, the functional group R is translocated to the monomer building block Initially carrying the nucleophile.

8

- (B) Nucleophilic attack by the amine on the thloester leads to formation of an amide bond, in effect translocating the functional group R of the thioester to the other monomer building block.
- (C) Reaction between hydrazine and p-ketoester leads to formation of pyrazolone, in effect translocating the R and R' functional groups to the other monomer building block.

22

(D) Reaction of hydroxylamine with p-ketoester leads to formation of the isoxazolone, thereby translocating the R and R' groups to the other

30 monomer building block.

PCT/DK2003/000417

2

(E) Reaction of thiourea with β-ketoester leads to formation of the pyrimidine, thereby translocating the R and R' groups to the other monomer building block.

- (F) Reaction of urea with malonate leads to formatlon of pyrimidine,
- thereby translocating the R group to the other monomer building

ß

(G) Depending on whether Z = O or Z = NH, a Heck reaction followed by a nucleophilic substitution leads to formation of coumarin or quinolinon, thereby translocating the R and R' groups to the other monomer build-

9

- (H) Reaction of hydrazine and phthallmides leads to formation of phthalhydrazide, thereby translocating the R and R' groups to the other monomer building block.
- (I) Reaction of amino acid esters leads to formation of diketopiperazine, thereby translocating the R group to the other monomer building

5

- (J) Reaction of urea with a-substituted esters leads to formation of hydantoin, and translocation of the R and R' groups to the other monomer building block.
- (K) Alkylation may be achieved by reaction of various nucleophiles with sulfonates. This translocates the functional groups R and R' to the other monomer building block.

2

(L) Reaction of a di-activated alkene containing an electron withdrawing and a leaving group, whereby the alkene is translocated to the nucleophile.

22

- (M) Reaction of disulfide with mercaptane leads to formation of a disulfide, thereby translocating the R' group to the other monomer building
- (N) Reaction of amino acid esters and amino ketones leads to formation of benzodiazepinone, thereby translocating the R group to the other monomer building block.

ဗ္က

WO 2004/001042

PCT/DK2003/000417

7

(O) Reaction of phosphonates with aidehydes or ketones leads to formation of substituted aikenes, thereby translocating the R" group to the other monomer building block.

- (P) Reaction of boronates with aryls or heteroaryls results in transfer of an aryl group to the other monomer building block (to form a blaryl).
  - (Q) Reaction arylsulfonates with boronates leads to transfer of the aryl

Ŋ

- group.
  (R) Reaction of boronates with vinyls (or alkynes) results in transfer of an
  - aryl group to the other monomer building block to form a vinylarene (or alkynylarene).

    (S) Reaction between aliphatic boronates and arylhalides, whereby the

9

- alkyl group is translocated to yleld an alkylarene.

  (T) Transition metal catalysed alpha-alkylation through reaction between an enolether and an arylhallide, thereby translocating the aliphatic
- (U) Condensations between e.g. enamines or enolethers with aldehydes leading to formation of alpha-hydroxy carbonyls or alpha,beta-unsaturated carbonyls. The reaction translocates the nucleophilic part.

5

(V) Alkylation of alkylhalldes by e.g. enamines or enolethers. The reaction translocates the nucleophillc part.

ឧ

- (W) [2+4] cycloadditions, translocating the diene-part.
- (X) [2+4] cycloadditions, transiocating the ene-part.
- (Y) [3+2] cycloadditions between azides and alkenes, leading to triazoles by translocation of the ene-part.
- (Z)·[3+2] cycloadditions between nitriloxides and alkenes, leading to Isoxazoles by translocation of the ene-part.

22

Fig. 5 shows collection of reactive groups that may be used for templated synthesis, along with the bonds formed upon their reaction. After reaction.

30 activation (cleavage) may be required.

22

Fig. 6 shows various cleavable linkers, the conditions for their cleavage, and the resulting products.

### Example

5 Example 1. An encoded molecule library of small molecules displayed on an array

This example describes how small molecules from an encoded library can be positioning on an array. These libraries are encoded by codons that codes for each chemical entity in the final displayed molecule. These codons describe the synthetic history which is directed by the template. This example will show how the codons can be used to positioning the displayed molecules on an array in a predetermining way trough the codon composition in each individual template molecule.

9

The example below shows the encoding process of a 27-membered library including the RGD (Arg-Gly-Asp) sequence. This tri-peptide sequence is known to bind to various integrins such as the  $\alpha_s \beta_3$  integrin, for example. The library has 27 different members because of the total amount of combinations possible for a tri-peptide (3³).

5

The scheme below shows the encoding of the RGD sequence as an example of all possible combination of these three different chemical entities. The annealing of the identifier (upper strand) to the building block (lower stand) allows the transfer of the chemical entities and their corresponding anti-codons to the nascent complex. The chemical entities are transferred by a chemical reaction and the information of the anti-codon are transferred to the nascent complex by extending the identifier using a polymerase and a mixture of dNTPs. The letters in bold indicate a flanking region of the anti-codon and the antitodon as well as the codon is underlined. I indicate the non-discriminating base inosine.

30

STEP 1A. Annealing

23

-D-CGT GTG ATC GAA CTC GTG TG GTATCCTGAGGGTAC

STEP 1B. Transfer of chemical entity D and anti-codon

-D-GCA CAC TAG CTT GAG CAC AC CATROGRACITOCCATO

5 -GCA CAC TAG CTT GAG CAC AC CAT<u>ICCTGA</u>GGGTAC

STEP 2A. Annealing

-D-GCA CAC TAG CTT GAG CAC AC CAPAGGACTCCCANG -Q-GGT GTG ATC GAA CTC GTG TG GTAIIIIIIIIIITACGAAGCTGTTACG

9

STEP 2B. Transfer of chemical entity G and anti-codon.

-0-D-GCA CAC TAG CTT GAG CAC AC CATAGGACTCCCAAGGCTCGACAATGC -CGF GTG ATC GAA CTC GTG TG GTAIIIIIIIIAAGGAAGGTGTTAGG

5

STEP 3A. Annealing
-q-p-can can take the canage constant constant con the canage con an economic seconds the canage canage canage con an economic seconds canage canage canage con an economic seconds canage con economic seconds canage con economic seconds canage con economic seconds canage con economic seconds

TOC COC -CGT GTG ATC GIA CTC GTG TG GTAIIIIIIIIIIIIIIIIIIIIIIIIGG<u>CTCTAG</u>GGGT GGA AAT GCT AGG GCG

22

The transfer of chemical entities is described in detail below. This scheme shows in the first step how D is transferred from a building block to the identifier molecule and then how G and R is transfer from the building block to the scaffold (an amine group). The final step is a deprotection to obtain the RGD peptide linked to the template. The final product is a library complexes,

30 peptide linked to the template. The final product is a library complexes, wherein each complex comprises a template and an encoded trl-peptide.

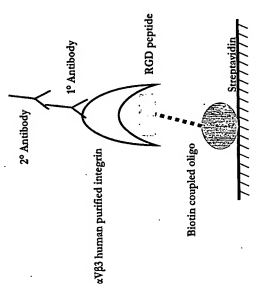
15

WO 2004/001042

25

The small molecule binding setup. In this example, the array setup was tested in a regular ELISA assay to confirm the binding of integrin to the immobilized RGD-template molecule. The ELISA was performed by immobilizing the template molecules using biotin binding to immobilized streptavidin.

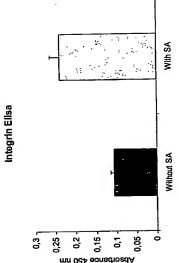
5 The component involved in this setup is shown below. The binding of integrin to the immobilized RGD-template molecule was detected using a specific antibody.



10

The result of the ELISA is shown below. The result shows that integrin binding is dependent of the immobilized RGD-template molecule. This setup with the immobilized RGD-template molecule is identical to the situation on the array except that the template molecule is immobilized through the probe that is complementary to the template molecule.

8



cat # P-9416), 3% casein (Sigma cat# C-8654) and 0,1 mg/ml herring sperm well strips (Pierce cat #15120) blocked over night in 0,5% Tween 20 (Sigma (Sigma ca拼 D-3159) In PBS . 3 pmol oligo/well diluted in 100 µl wash buffer 0,5% Tween 20, 3% casein In PBS) 1 hour shaking at r.t. After washing 5 The ELISA was performed using a 20-mer 5' biotinylated and 3' cRGD peptide coupled oligos This construct were bound to precoated streptavidin 8 limes, block buffer was added and incubated another hour at r.t shaking.

body was detected by TMB plus substrate (Kem-En-Tech cat #4390L), 100 µl 111: 2159-2170) (Chemicon cat # CC1018) was added to each well in 100 µl vash buffer containing 1mM MnCl<sub>2</sub> and incubated at room temperature shak-Jase conjugated antibody (abcam cat # ab6728) incubation, 100 µl 1:2000 Ing for 2 hours. After washing away unbound integrin,  $\alpha_V\beta_3$  Integrin mouse dilutions in wash-buffer, 1 hour shaking at r.t. After washing bound 2° anti-0,1µg/well αvβ3 human purified integrin (Belkin V.M et al (1990) J.Cell boli. (Martin-Padura I., et al (1995) J. Path. 175: 51) was added 0,05 µg/well In washes and subsequently 2° polyclonal to mouse IgG horseradish peroximonoclonal primary antibody (abcam cat # Ab7167) previously described 100 µl wash buffer, incubated shaking at r.t for 1 hour. Followed by 10

5

9

added and incubated until sufficient color development, to which 100 µl 0,2M

೪

corded at 450 nm. Control used for the experiment was a none binding com-H<sub>2</sub>SO<sub>4</sub> was added to stop the reaction. Absorbance was measured and remonly used RAD peptide coupled to an oligo (J Control Release 2002 Oct 4;83(2):241-51).

complementary to all the possible variations of codons on the Identifier molecule as well as a flanking sequence showing the position of the codons. The igure below shows the outline of the setup with the adaptor oligo binding to will display the motecule on the array determined by the codons in the temboth the identifier molecule as well as to the probe oligo on the array. This Identification of encoded library members using arrays. GenFlex from Affymetrix is used to designing so called adaptor oligos which are complementary to the different probe oligos on the array. The adaptor oligos are also 9

e.q. RGD

5

Adaptor

The generation of an encoded library gives a theoretical possibility of 27 difsigned for which amino acid it is encoding. Flanking sequences of three nuvariations generated by utilizing DNA array; codons were as described deferent variants. To efficlently enable identification of all possible tripeptide

8

PCT/DK2003/000417

cleotides around every peptide codon was added to ensure the precise binding between the template and adaptor molecule.

In order to detect all possible variants in this RGD library, 27 different adaptor oligos has to be designed that recognizes each Individual template molecule.

ß

bold letters is the complementary sequence to the various probes on the chip from Affymetrix. This chip contains 2000 different probes, of 20-mers bound therefore permit the binding of the template-displayed molecules. The adapto the chip. These 27 different adaptor oligos are shown below, where the and the normal letter sequence is the part that corresponds to the codons tor oligos were designed to be complementary to probes on the GenFlex These adaptor oligos will bind specifically to the probes on the array and and flanking sequences described above.

9

RCCTCTAGTG 5

è

'n

- CATANGETT TANGETHERE TO THE WANTERNIGON ೪

CATCCTCTAGTGATGCCTCTAGTGTGCCCTCTAGTGGCA

- CATRICCTICA GGGATGTCCTCA GGGTGCTTGGTGAGGGGCA
- CATCCTCTAGTGATGGGTATTGCTTTTGGTATTGGA CATCCTCTAGTGATGCCTCTAGTGTGCTARY SIMIGCA
  - CATCCTCTAGTGATGTCTGHGGGTGCTTGTTGTTGTGGGCCA
- CATCCTCTAGTGATGCCTCTAGTGTGGTGTGTGGTGGTGCCA 22
  - CATOMINERIALISCCICIAGICCCCCCCTCTAGICCCA
- CATE HERET AND THE STREET TOCCCTCTAGTGGCA 9
  - CATHCCTGAGGGATCHANGARTTGCCCTCTAGTGGCA CATTCCTGAGGGATGCCTCTAGTGTGCCCTCTAGTGGCA Ξ. 12. ဓ္က
- 13. CATURAGO CONTRACTO CIGAGO GIGO COLOTA GIGO CA CATERER PROBLEM TOTCCTONGGGTGCTCCTGAGGGGGCA
- CATELLEGIZATCHEREREFIECTICCTGAGGGGCA

WO 2004/001042

PCT/DK2003/000417

16. carrecreacegaregraver - recreereaced

CATCCTCTAGTGATGWNPSNY TTGCTCCTGAGGGGCA

19 CATICETERGEGATGRANS (A) TGCCCTCTAGTGGCA

CATHOCTIONGGGATGTCCTIGAGGGTGCCCTCTAGTGGCA 

S

CATCCTCTAGTGATGTCCTGAGGGTGCTHY 22.

24. CATCCTCTAGTGATGATATE TGCCCTCTAGTGGCA 23. CATE NATIONAL PROCESSOR OF CONTRACT AND THE CONTRACT OF TH

CATALLANT ATCHERENCE TO CATALLY TO THE CAL CATTECTICAGGGATGCCTCTAGTGTGTTCTTGAGGGGCA

5

CATCCTCTAGTGATGTGAGGGTGCCCTCTAGTGGCA

combinations (Marked in bold, complementary sequence to probe sequence Adaptor oligos to be used for DNA array analysis for all RGD-library codon

with the probe set number on the chip to the right).

5

Sequence 5'-3'Probe set # on chip Primer#

೪

1. GCT AGG CTA ATG TCC GGC TAG TAG GAG ATCA CTA CGG AGA TCA CAC GGG AGA TCA CC GT 90109

2. AGG CAG ACA ACT CAA TCC GGG TAC TTC GAC AAT ACC TTC GAC AAA GGC TTC GAC AAC GT

22

3. TCA GAC TAG GGT AGG GGA TAG TAA GGA CTC CCT ACA GGA CTC CCA CGA GGA CTC CCC GT

4. TGT CCA GTA GCT TGA GAG TCG TAG QAG ATC ACT ACC TTC GAC AAA CGC TTC GAC AAC GT

ဓ္က

00445

5. CGA CAA GGC ATT CAC ACT AGG TAG GAG ATC ACT ACG GAG ATC ACA CGC TTC GAC AAC GT

38

8. TCG GCG TTA CGT GCT GAC TAG TAG GAG ATC ACT ACA GGA CTC CCA CGA GGA CTC CCC GT

7. CCG CAG CAA GCT ATC QAG AAG TAG QAG ATC ACT ACG GAG ATC ACA CGA GGA CTC CCC GT

<del></del>

WO 2004/001042

9, GTA CGT TGA CAG TCT GCA CAG TAC TTC GAC AAT ACG GAG ATC ACA CGG GAG ATC ACC GT IDDES2

S

10, GTC TGG CCC TAC CTA TGG TTG TAC TTC GAC AAT ACC TTC GAC AAA CGG GAG ATC ACC GT

10 11. ACE ANT GCA ANT AGG CGG CCG TAA GGA CTC CCT ACC TTC GAC AAA CGG GAG ATC ACC GT 00922

12. TCA GGC CCA CGT AGC GTT ATG TAA GGA CTC CCT ACG GAG ATG ACA CGG GAG ATC ACC GT

13. GAA CTA TGC TGA CAG TAC CGG TAC TTC GAC AAT ACA GGA CTC CCA CGG GAG ATC ACC GT 01081

5

14. CCC AGG GCA AGG GAT GAT AAG TAC TTC GAC AAT ACA GGA GTG CCA CGA GGA CTC CCC GT

8

15, TCA CGT AAT TTG TTA GCC GCG TAC TTC GAC AAT ACC TTC GAC AAA CGA GGA CTC CCC GT

18. TAC CTG GCA TGA CGC GAT ATG TAA GGA CTC CCT ACC TTC GAC AAA GGA GGA CTC CCC GT

22

17. TGC AGG CTC GCA GAT GCT ATG TAA GGA CTC CCT ACA GGA CTC CCA CGC TTC GAC AAC GT M414 18. TOA GGO TTA GAG CTT GAT CGG TAG GAG ATC ACT ACC TTG GAC AAA CGA GGA CTC CCC GT 01622

ဗ္က

18, TET CGG TTA VTG AGT GGA CTG TAA GGA CTC CCT ACC TTC GAC AAA 09G GAG ATC ACC GT 01528 20, CGA CGA GCA CCA ATT CGA GAG TAA GGA CTC CCT ACC TTC GAC AAA CGC TTC GAC AAC GT

32

40 21.9TT AGA TCA TGA TCG CGG TGG TTA GGA CTC CCT ACA GGA CGC CCA CGG GAG ATC ACC GT

22. CAC TAA GAC ATG CAC AGG GGG TAG GAG ATC ACT ACA GGA CTC CCA CGC TTC GAC AAC GT

23. CTA CTG ACA CTG ACC AGG GAG TAC TTC GAC AAT ACG GAG ATC ACA CGC TTC GAC AAC GT DIRID

45

3

24. GCA TAC AGG TTA CGA CGC CTG TAG GAG ATC ACT ACC TTC GAC AAA CGG GAG ATC ACC GT n1882 28. CTT CGC GCA GCT ACA TAS ATG TAA GGA CTC CCT ACG GAG ATC ACA CGA GGA CTC CCC GT

S

01828 28. GGC ATA CTA GAG TCA GGG ATG TAC TTC GAC AAT ACA GGA CTC CCA CGC TTC GAC AAC GT 10 27. ATC AAG GCA ACC GCC ABT AGG TAG GAC ATC ACT ACA GGA CTC CCA CGG GAG ATC ACC GT

15

The adaptor oligos and the templates with displayed molecules were used for analysis on GenFlex according to protocol below.

GenFlex hybridisation and scanning. Prior to hybridization, the Adaptor mix (100 nM each final concentration) in a hybridization buffer (100mM MES, 1 M

ន

(100 pM each final concentration) in a hybridization buffer (100mM MES, 1 M NaCl, 20 mM EDTA, 0,01% Tween 20, 1x Denhardt's), was heated to 95°C for 5 min and subsequently to 40°C for 5 min before loading onto the Affymetrix GenFlex probe array cartridge. The probe array was then incubated for 2h at 45°C at constant rotation (60 rpm). The Adaptor mix was removed from the GenFlex cartridge, and replaced with the Template in a hybridization buffer

33

(100mM MES, 1 M NaCl, 20 mM EDTA, 0,01% Tween 20, 1x Denhardt's). The Template hybridisation mix was heated to 95°C for 5 min and subsequently to 40°C for 5 min before loading onto the Affymetrix GenFlex probe array cartridge and hybridised for 2h at 45°C at constant rotation (60 rpm). The washing and staining procedure was performed in the Affymetrix Fluidics Station. The probe array was exposed to 2 wash in 6xSSPE-T at 25°C followed by 12 washes in 0.5xSSPE-T at 40°C. Then, 0,1µg/well ανβ3 human purified in-

ဗ္က

CC1018) was added to the chip in wash buffer containing1mM MnCl<sub>2</sub> and incubated at room temperature shaking for 2 hours. After washing away unbound integrin, α<sub>Vβ3</sub> integrin mouse monoclonal primary antibody (abcam cat

tegrin (Belkin V.M et al (1990) J.Cell boil. 111: 2159-2170) (Chemicon cat #

PCT/DK2003/000417

32

# Ab7167) previously described (Martin-Padura I., et al (1995) J. Path. 175: 51) was added in wash buffer, and rotated at room temperature for 1 hour. Then, after 5-10 washes a 2° polyclonal to mouse IgG horseradish peroxidase conjugated antibody (abcam cat # ab6728) was incubation in the washbuffer. After washing off non-bound 2° antibody the probe arrays were scanned at 560 nm using a confocal laser-scanning microscope with an argon ion laser as the excitation source (Hewlett Packard GeneArray Scanner

S

WO 2004/001042

PCT/DK2003/000417

33

### Claims

- A microarray comprising a plurality of single stranded nucleic acid probes immobilized in discrete areas of a solid support, said probes being
- hybridised to a library of complexes, wherein each complex comprises an encoded molecule and a template which codes for said molecule, said template comprising a number of codons which codes for chemical entities which upon reaction form a reaction product which at least partly form part of the encoded molecule.
- A microarray according to claim 1, wherein the chemical entities are precursors for a structural unit appearing in the encoded molecule.
- A microarray according to claim 1 or 2, wherein the chemical entities are transferred to the nascent encoded molecule by a building block, which further comprises an anti-codon.
- A microarray according to claim 3, wherein the information of the anticodon is transferred in conjunction with the chemical entity to the nascent complex.
- A microarray according to any of the claims 1 to 4, wherein the chemical entities are reacted without enzymatic interaction.
- 6. A microarray according to any of the claims 1 to 5, wherein the template comprises two or more codons.
- 7: A microarray according to any of the claims 1 to 6, wherein the nucleic acid probe of the array is hybridised to a template through an adapter oligonucleotide having a sequence complementing the probe as well as
- 25 the template.
- A method for preparing a microarray displaying a library of encoded molecules, wherein an oligonucleotide microarray comprising a plurality of single stranded nucleic acid probes immobilized in discrete areas of a solid support is mixed under conditions which allows for specific
- hybridisation with a library of complexes, each of sald complexes comprising an encoded molecule and a template which codes for sald molecule, sald template comprising a number of codons which codes for

ജ

34

chemical entities which upon reaction form a reaction product which at

least partly form part of the encoded molecule.

A method for identifying an encoded molecule having a preselected property, comprising the steps of i) providing the microarray according to claim 1

σ.

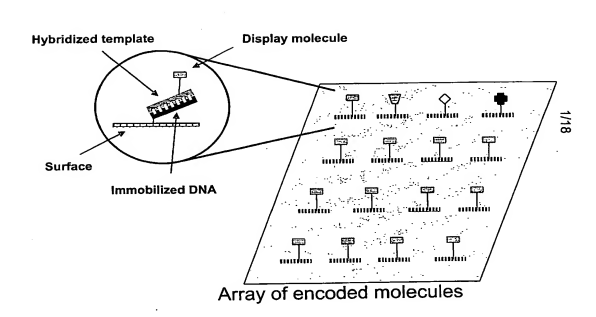
iii) washing non-bound material off, and iv) detection any hound material in each specific

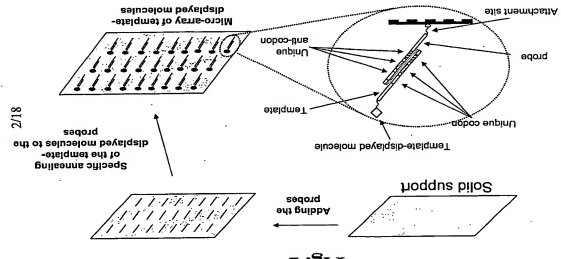
ii) adding a biological sample containing target molecules,

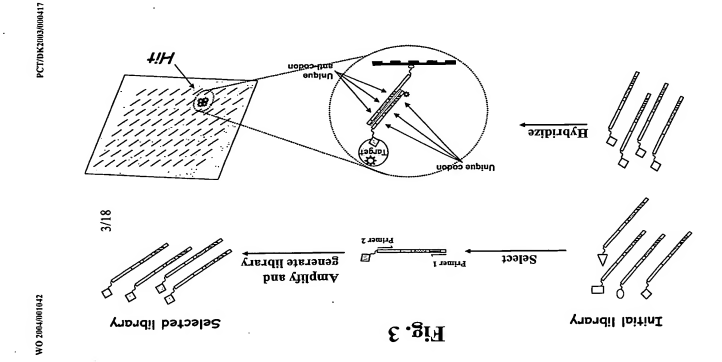
- iv) detecting any bound material in each spot.
- 10. Use of a microarray according to any of the claims 1 to 7 for identifying an encoded molecule capable of binding to a target molecule.

ಕ

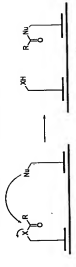
Fig. 1





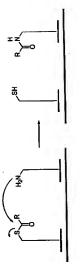


A. Acylating monomer building blocks - principle



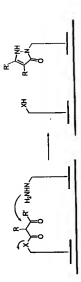
B. Acylation

Amide formation by reaction of amines with activated esters



C. Acylation

Pyrazolone formation by reaction of hydrazines with  $\beta\text{--}Ketoesters$ 



5/18

PCT/DK2003/000417

Isoxazolone formation by reaction of hydroxylamines with

β-Ketoesters

E. Acylation

Pyrimidine formation by reaction of thioureas with  $\beta\text{--}Ketoesters$ 

F. Acylation

Pyrimidine formation by reaction of ureas with Malonates

6/18

### G. Acylation

Coumarine or quinolinon formation by a Heck reaction followed by a nucleophilic substitution

### H. Acylation

Phthalhydrazide formation by reaction of Hydrazines and Phthalimides

### I. Acylation

Diketopiperazine formation by reaction of Amino Acid Esters

WO 2004/001042

CONTURNE

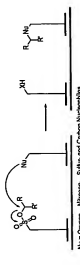
7/18

### J. Acylation

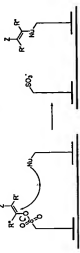
Hydantoin formation by reaction of Urea and  $\alpha\text{-substituted}$  Esters

## K. Alkylating monomer building blocks - principle

Alkylated compounds by reaction of Sulfonates with Nucleofiles



## L. Vinylating monomer building blocks - principle



Z = CN, COOR, COR, NO<sub>2</sub>, SO<sub>2</sub>R, S(=O)R, SO<sub>2</sub>NR<sub>2</sub>, F No a Occasio, Nilmona, Sulfar, and Carbon Markemblia

M. Heteroatom electrophiles

Disulfide formation by reaction of Pyridyl disulfide with Mercaptanes

### N. Acylation

Benzodiazepinone formation by reaction of Amino Acid Esters and Amino Ketones

WO 2004/001042

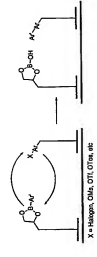
9/18

## O. Wittig/Horner-Wittig-Emmons reagents

Substituted alkene formation by reaction of Phosphonates with Aldehydes or Ketones

### P. Arylation

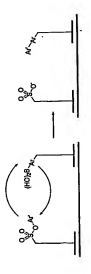
Biaryl formation by the reaction of Boronates with Aryls or Heteroaryls



10/18

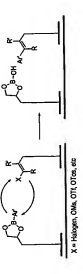
Q. Arylation

Blaryl formation by the reaction of Boronates with Aryls or Heteroaryls



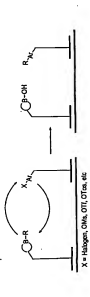
R. Arylation

Vinylarene formation by the reaction of alkenes with Aryls or Heteroaryls



S. Alkylation

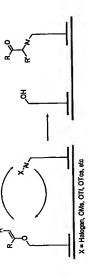
Alkylation of arenes/hetarens by the reaction with Alkyl boronates



41/18

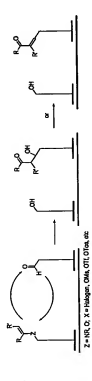
T. Alkylation

Alkylation of arenas/hetarenes by reaction with enolethers



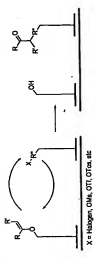
U. Condensations

Alkylation of aldehydes with enolethers or enamines



V. Alkylation

Alkylation of aliphatic halides or tosylates with enolethers or enamines



12/18

W. [2+4] Cycloadditlons

X. [2+4] Cycloadditions

Y. [3+2] Cycloadditions

13/18

Z. [3+2] Cycloadditions

Fig. 5

14/18

### . Nucleophilic substitution reaction

ZZ = COOR, CHD, COH, CONF2, COO',
NO, SOR, SO2R, SO2HF3, CM, ed.

## Aromatic nucleophilic substitution

Transition metal catalysed reactions

SUBSTITUTED AROMATIC COMPOUNDS

RETORNE REPORT RECOMPOUNDS

RETORNE RESTRUCTOR ARCHANTIC COMPOUNDS

RETORNE RESTRUCTOR RETORNE RETORNE RESTRUCTOR RETORNER RETORN

15/18

## Addition to carbon-carbon multiplebonds

## Cycloaddition to multiple bounds

16/18

## Addition to carbon-hetero multiple bonds

17/18

Fig. 6

A. Linker for the formation of Ketones, Aldehydes, Amides and Acids

B. Linker for the formation of Ketones, Amides and Acids

C. Linker for the formation of Aldehydes and Ketones

D. Linker for the formation of Alcohols and Acida

E. Linker for the formation of Amines and Alcohols

F. Linker for the formation of Esters, Thioesters, Amides, and Alcohols R. O. H. T. W. T. O.H. O. H. J. W.

G. Unker for the formation of Sulfonamides and Alcohols

PCT/DK2003/000417

18/18

H. Linker for the formation of Ketones, Amines and Alcohols

. Linker for the formation of Ketones, Amines, Alcohols and Mercaptanes

J. Linker for the formation of Blaryl and Bihetaryl

K. Linker for the formation of Benzyles, Amines, Anilins Alcohols and Phenoles

L. Linker for the formation of Mercaptanes

M. Linker for the formation of Glycoside:

POS NOS THE POST OF THE POST O

N. Linker for the formation of Aldehydes and Glyoxylamides HA CHAIG. HAS HE SHIP TO BE

O. Linker for the formation of Aldehydes, Ketones and

# (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property International Bureau Organization



(10) International Publication Number WO 2004/001042 A3

PCT

Per-Ola [SPSRI]; Ringvilgen 7, S-235 93 Veilingo (SP). PEDENSEN, Henrik [DK/DK]; Prodesvej 24, DK-2880 Bagsværd (DK).

C12Q 1/68, 31 December 2003 (31.12.2003)

(43) International Publication Date

(51) International Patent Classification?: C12N 15/10 // C12P 21/02, C07II 21/00

PCT/DK2003/000417 (21) International Application Number:

(22) International Filing Date: 20 June 2003 (20.06.2003)

(25) Filing Language:

English

(26) Publication Language:

Hnglish

7 (81) Designated States (national): AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DH, DB, BD, ED, EB, EB, BG, BB, GD, GR, GH, CM, IR, HU, LD, LL, N, SL, PK RK, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MY, MY, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SI, SG, SK, SI, TY, TM, TN, TR, TT, ZU, UG, US, UZ, Mh VC, VN, YU, ZA, ZM, ZW.

Designated States (regional): ARIPO patent (GH, GM, KH, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurusian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), Buropean patent (AT, BH, BG, CH, CY, CZ, DE, DK, HH, IS, Ft, FR, GB, GR, HU, HE, TT, LU, MC, NI., FT, RO, SE, SI, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NB, SN, TD, TG). 3

2 2 2 2 S

20 June 2002 (20:06:2002) 20 June 2002 (20:06:2002)

PCT/DK02/00419

(30) Priority Data: 10/175,539 50/434,664

20 December 2002 (20.12.2002)

PA 2002 01953 19 December 2002 (19.12.2002)

Published:

(71) Applicant (for all designated States except US); NUEVO-LUTION A/S [DK/DK]; Rønnegade 8, 5, DK-2100

with international scarch report before the expiration of the time limit for amending the claims and to be republished in the event of revelpt of

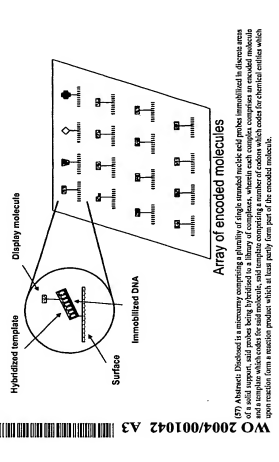
[Continued on next page]

## (54) THE: MICROARRAYS DISPLAYING ENCODED MOLECULES

(75) Inventors/Applicants (for US only): FRESKGÅRD,

(72) Inventors; and

København Ø (DK).



# WO 2004/001042 A3 INCOMENIA INCOMENI

(88) Date of publication of the international search report: 18 March 2004

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazene.

## INTERNATIONAL SEARCH REPORT

A CLASSIFICATION OF BUBLICT MATTER //C12P21/02, C07H21/00 IPC 7 C12Q1/68 C12N15/10 //C12P21/02, C07H21/00

In at Application No PCT/DK 03/00417

Relevant to dalm No. The later document published after the interrational filing date or printing that and not incentified, which the supplication but also also also the contification but also the control of 1-10 1-10 2 Date of mailing of the International search report Patent family members are listed in ermex. Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched 'à' document member of the same patent tenuly Electronic data base consulted during the international search (name of data base and, where practical, search forms used) Osborne, H 10/02/2004 Authorized officer SUMMERER D-ET AL: "DNA-TEMPLATED SYNTHESIS:\*MORE #ERSATILE THAN EXPECTED" ANGEMANDTE. CHEMIE. INTERNATIONAL EDITION, YERAGE CHEMIE. WEINHEIM, DE, vol. 41, no. 1, 40 anuary 2002. (2002-01-04), pages 89-90, XP001170352 Category\* Citation of document, with Indication, where appropriate, of the relevant passages According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED

Minimum documentation searched (clear/fiction system followed by disselfcation symbols)

IPC 7 C12Q C12N C07H C12P US 5 723 598 A (BRENNER SYDNEY ET AL) 3 March 1998 (1998-03-03) column 1 -column 3, line l <del>|</del> WO 00 23458 A.(UNIV LELAND STANFORD JUNIOR) 27 April 2000 (2000-04-27) page 1 -page 3, "ilne 30 EPO-Internal, MEDLINE, BIOSIS, EMBASE X Further documents are listed in the continuation of box C. Nume and moling address of the ISA
European Patent Offea, P.3. 5618 Patentlaan 2
II. – Zoot HV Tigwiti
Tal (-51.–70) 340–3016
Fex (-51.–70) 340–3016 "P document published prior to the international filing date but later than the priority date delimed \*E\* earlier document but published on or after the International filing date Cournett which may laww debtas on profity talling) or which is dead to elabytism he publication date of another with or other apostal reason (as apostiles).
 Courner referring to an oral deadesture, use, antibition or other means. A document defining the general state of the art which is not considered to be of particular relevance. Date of the solual completion of the International search C. DOCUMENTS CONSIDERED TO BE RELEVANT ISSN: 0570-0833 the whole domument Special categories of citod documents: 3 February 2004

Form PCT/ISA210 (second sheet) (July 1992)

	INTERNATIONAL SEARCH REPORT	Int ional Application No PCT/DK 03/00417	100 No
Continu	C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Ctation of document, with indication, where appropriate, of the relevant passages	Refe	Relevant to claim No.
>-	BRUICK R K ET AL: "TEMPLATE-DIRECTED LIGATION OF PEPTIDES TO OLIGONUCLEOTIDES" CHEMISTRY AND BIOLOGY, CURRENT BIOLOGY, LONDON, 6B, vol. 3, no. 1, January 1996 (1996-01), pages 49-56, XPO00856876 ISSN: 1074-5521 the whole document		1-10
<b>&gt;</b> -	WALDER J A ET AL: "COMPLEMENTARY CARRIER FEFTIDE SYNTHESIS: GENERAL STRATEGY AND IMPLICATIONS FOR PREBIOTIC ORIGIN OF PEPTIDE SYNTHESIS" PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCE. OF USA, NATIONAL ACADEMY OF SCIENCE. MASHINGTON, US, VOl. 76, no. 1, January 1979 (1979–01), pages 51–55, XPO00857351 ISSN: 0027–8424 the whole document		1-10
<b>&gt;</b> -	EP 0 778 280 A (ISIS INNOVATION) 11 June 1997 (1997-06-11) figures 4,5; example 17		1-10
<b>&gt;</b> -	E WO 96 41011 A (SPECTRAGEN INC) - 19 December 1996 (1996-12-19) 7 figure 3		1-10
<b>&gt;-</b>	LOCKHART D J ET AL: "EXPRESSION MONITORING BY HYBRIDIZATION TO HIGH-DENSITY OLIGONUCLEOTIDE ARRAYS" BIO/TECHNOLOGY, NATURE PUBLISHING CO. NEW YORK, US, vol. 14, no. 13, vol. 14, no. 13, vol. 10 December 1996 (1996-12-01), pages 1675-1680, XP002022521 ISSN: 0733-222X the whole document		1-10
<b>&gt;</b>	WO 97 27317 A (CHEE MARK ;LAI CHAOQIANG (US); LEE DANNY (US); AFFYMETRIX INC (US)) 31 July 1997 (1997-07-31) figures 12-15		1-10
<b>&gt;-</b>	WO 99 42605 A (SCHWABACHER ALAN W) 26 August 1999 (1999-08-26) page 2, 11ne 11 - 11ne 16		1-10
⋖	WO 94 08051 A (READER JOHN C ;STILL W CLARK (US); UNIV COLUMBIA (US); COLD SPRING) 14 April 1994 (1994-04-14)		

$\vdash$
ì
ਰ
盃
ш
~
=
六
$\approx$
5
9
Ж.
0,
٦
⊴
Ž
Ō
F
⋖
Z
œ
ш
⊢
Z

mormation on pattern taitary memores

Intu mal Application No PCT/DK 03/00417

Publication date	08-05-2000 27-04-2000 16-08-2001 27-04-2000	12-11-1996 09-05-2000 15-06-2000 15-01-1998 08-11-1993 14-10-1993 06-07-2000 26-04-2001 07-08-2000 22-03-1995 01-09-2000 22-06-1995	11-06-1997 15-11-1997 15-11-1997 13-08-1998 28-02-1995 06-02-1996 06-02-2003 22-12-1997 15-02-1996 16-12-1997 29-01-1996 09-02-1995 16-10-1996 10-06-2002 25-02-1997 28-03-1996 27-10-2000 22-08-2002 23-06-1998 18-10-2001	03-06-1997 13-04-2000 30-12-1996 19-12-1996 16-09-1998 17-06-1998 01-04-1999 06-07-1999 06-07-1999 05-02-1998 19-12-1996 18-11-1999
Patent tamily momber(s)	1318400 A 2346989 A1 1123305 A1 0023458 A1	5573905 A 6060596 A 193561 T 685050 B2 3944933 A 2132103 A 69328781 D1 69328781 T2 643778 T3 064377 T3 750530 T	0778280 A2 159767 T 230409 T 695349 B2 7259194 A 2168010 A1 1131440 A , B 69406544 D1 69406544 D1 7711362 T3 0711362 T3 0711362 A1 2108479 T3 960403 A 73802 A2 73802 A2 2269911 B2 9601830 C2 200115091 A1 60370 A 2002116091 A1 60370 A 200115091 A1 60370 A	5635400 A 718357 B2 6102096 A 2222581 A1 2332731 A1 1193357 A 9703926 A3 0832287 A1 9900910 A2 11507528 T 975744 A 324000 A1 9641011 A1
	8558	M SE E E E E E E E E E E E E E E E E E E	S C C C C C C C C C C C C C C C C C C C	AU PUP ECCA AUS AUS AUS AUS AUS AUS AUS AUS AUS AU
Publication date	27-04-2000	03-03-1998	11-06-1997	19-12-1996
	A	₹ .	<b>∢</b>	⋖
Patent document cited in search report	MO 0023458	US 5723598	EP 0778280	W0 9641011

Form PCT/SSAZ10 (patient family armod (July 1992)

INTERNATIONAL SEARCH REPORT Intermediation on patent family members

inte not Application No PCT/DK 03/00417

	1.			
Publication date	06-05-1996 30-04-1997 25-04-1997 10-09-1997 10-09-1997 28-07-1998 02-06-1997 28-08-2001 17-04-1997 22-05-2001	20-08-1997 02-12-1998 28-05-2002 31-07-1997 25-09-2003 03-04-2003 05-02-2002	17-01-2002 06-09-1999 26-08-1999 06-12-2000 26-11-2002 26-08-1999	15-07-2003 02-03-2000 12-02-1998 26-04-1994 14-04-1994 14-04-1995 28-05-1996 31-10-2003 02-07-1996 31-10-1996 14-04-1998 04-08-1998 04-08-1998 04-08-1998
Patent family member(s)	4277896 A 7717596 A 2202167 A1 9700866 A3 0793718 A1 0931165 A1 971473 A 10507357 T 971644 A 6280935 B1 9713877 A1 6235475 A1	2253397 A 0880598 A1 2002515738 T 9727317 A1 2003160757 A1 2003064364 A1 6344316 B1 2002165372 A1	2002006604 A1 2871299 A 2321111 A1 1056822 A1 2002540380 T 9942605 A1	244769 T 716621 B2 4525897 A 685579 B2 553694 A 2243848 A1 69333087 D1 665897 T3 0665897 A1 72456 A2 107166 A 8506.75 T 951230 A 9408051 A1 5565334 A 5789172 A 6503759 B1
Publication date	A A A A A A A A A A A A A A A A A A A	31-07-1997 AU SP SP S	26-08-1999 US AU CA CA EP DP UNO	14-04-1994 AU
Patent document clted in search report	WO 9641011 A	WO 9727317 A	MO 9942605 A	MD 9408051 A